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## **CLAIMS**

## WHAT IS CLAIMED IS:

- 1. A method for determining a preferred alignment between a query sequence and at least one template sequence comprising the steps of:
- 5 a. aligning at least two reference sequences to determine one or more BRIDGE/BULGE gaps;
  - b. determining at least one alignment score between the potential alignment of said query sequence and each said template sequence; wherein each said alignment score reflects whether said alignment between said query sequence and each said template sequence creates a BRIDGE/BULGE gap; and
  - c. determining a preferred alignment between said query sequence and each said template sequence based on said alignment score.
  - 2. The method of claim 1 wherein said preferred alignment is the optimal alignment.
  - 3. The method of claim 1 wherein step b comprises the steps of:
    - a. forming a sequence alignment similarity matrix for said query sequence and each said template sequence with matrix elements  $\mathbf{s}_{ij}$ ; and
    - b. determining a sequence alignment sum matrix with matrix elements  $S_{ij}$  from the dynamic evolution of each said sequence alignment similarity matrix, wherein the matrix elements of each said sum matrix reflect whether or not any potential alignment gaps that may be formed from the alignment of said query sequence with each said template sequence creates a BRIDGE/BULGE gap.
  - 4. The method of claim 3 wherein step b comprises the steps of:

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- a. calculating said sequence alignment sum matrix from the dynamic evolution of each said sequence alignment similarity matrix, according to the equation:
- $S_{ij} = s_{ij} + Max\{S_{i+1,j+1}, S_{i+1, j+2 \text{ to } jmax} GAP, S_{i+2 \text{ to } imax, j+2} GAP, S_{m,n} BRIDGE/BULGE\},$

wherein GAP represents the gap penalty for an alignment gap between said query sequence and each said template sequence, BRIDGE/BULGE represents the penalty for a known bridge or bulge that begins at the m,n matrix element of said sum matrix and ends at the i,j matrix element of said sum matrix and Max $\{S_{i+1,j+1}, S_{i+1,j+2 \text{ to jmax}} - GAP, S_{i+2 \text{ to imax}, j+2} - GAP, S_{m,n} - BRIDGE/BULGE\}$  refers to the maximum value of the four terms contained within the brackets

- 5. A method for determining a preferred alignment between a query sequence and at least one template sequence comprising the steps of:
  - a. aligning at least two reference sequences to determine at least one
    BRIDGE/BULGE gap;
  - b. forming a sequence alignment similarity matrix from said query sequence and each said template sequence;
- c. determining a sequence alignment sum matrix from the dynamic evolution of each said sequence alignment similarity matrix, wherein the matrix elements of each said sum matrix reflect whether or not any potential alignment gaps that may

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be formed from the alignment of said query sequence with each said template sequence creates a BRIDGE/BULGE gap; and

- d. determining a preferred alignment between said query sequence and each said template sequence from said dynamic evolution of each said sum matrix.
- 5 6. The method of claim 5 wherein said preferred alignment is the optimal alignment.
  - 7. A method for determining a preferred alignment between a query sequence and at least one template sequence comprising the steps of:
    - a. aligning at least two reference sequences to determine at least one BRIDGE/BULGE gap;
    - b. calculating a sequence alignment similarity matrix with matrix elements  $s_{ij}$  for said query sequence and each said template sequence;
    - c. calculating a sequence alignment sum matrix with matrix elements  $S_{ij}$  from the dynamic evolution of each said sequence alignment similarity matrix, according to the equation:

$$\begin{split} S_{ij} &= s_{ij} + Max\{S_{i+1,j+1},\, Si_{+1,\,j+2\,\,to\,\,jmax} - GAP,\, S_{i+2\,\,to\,\,imax,\,j+2} - GAP,\, S_{m,n} - \\ BRIDGE/BULGE\}, \end{split}$$

wherein GAP represents the gap penalty for an alignment gap between said query sequence and each said template sequence, BRIDGE/BULGE represents the penalty for a known bridge or bulge that begins at the m,n matrix element of said sum matrix and ends at the i,j matrix element of said sum matrix and Max $\{S_{i+1,j+1},$ 

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 $S_{i+1, j+2 \text{ to } jmax} - GAP$ ,  $S_{i+2 \text{ to } imax, j+2} - GAP$ ,  $S_{m,n} - BRIDGE/BULGE$ } refers to the maximum value of the four terms contained within the brackets; and

- d. determining a preferred alignment between said query sequence and each said template sequence from the dynamic evolution of said sum matrix.
- 5 8. The method of claim 7 wherein said preferred alignment is an optimal alignment.
  - 9. A method for determining a preferred alignment between at least one query sequence and at least one template sequence, for use in primary sequence homology modeling methods, comprising the steps of:
    - a. aligning at least two reference sequences to determine one or more
      BRIDGE/BULGE gaps;
    - b. determining at least one alignment score for the potential alignment of each said query sequence and each said template sequence; wherein each said alignment score reflects whether said alignment between each aid query sequence and each said template sequence creates a BRIDGE/BULGE gap; and
    - c. determining a preferred alignment between each said query sequence and each said template sequence based on said alignment scores, wherein said preferred alignment contains from approximately 10% to approximately 20% homologous residues.
  - 10. The method of claim 9 wherein said preferred alignment is the optimal alignment.
- 20 11. The method of claim 9 wherein said primary sequence homology method is a method for determining the three dimensional structure of said query sequence.

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- 12. The method of claim 10 wherein said primary sequence homology modeling method is a method for determining the three dimensional structure of said query sequence.
- The method of claim 9 wherein said primary sequence homology modeling
  method is a method for determining the primary sequence homology relationship
  between at least two query sequences.
  - 14. The method of claim 10 wherein said primary sequence homology modeling method is a method for determining the sequence homology relationship between at least two query sequences.
- 10 15. A method for determining a preferred alignment between at least one query sequence and at least one template sequence, for use in primary sequence homology modeling methods, comprising the steps of:
  - a. aligning at least two reference sequences to determine at least one
    BRIDGE/BULGE gap;
  - b. forming a sequence alignment similarity matrix for each said query sequence and each said template sequence;
  - c. determining a sequence alignment sum matrix from the dynamic evolution of each said sequence alignment similarity matrix, wherein the matrix elements of each said sum matrix reflect whether or not any potential alignment gaps that may be formed from the alignment of each said query sequence with each said template sequence creates a BRIDGE/BULGE gap; and
  - d. determining a preferred alignment between each said query sequence and each said template sequence from said dynamic evolution of each said sum

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matrix, wherein said preferred alignment contains from approximately 10% to approximately 20% homologous residues.

- 16. The method of claim 15 wherein said preferred alignment is the optimal alignment.
- 5 17. The method of claim 15 wherein said primary sequence homology method is a method for determining the three dimensional structure of said query sequence.
  - 18. The method of claim 16 wherein said primary sequence homology modeling method is a method for determining the three dimensional structure of said query sequence.
- 10 19. The method of claim 15 wherein said primary sequence homology modeling method is a method for determining the primary sequence homology relationship between at least two query sequences.
  - 20. The method of claim 16 wherein said primary sequence homology modeling method is a method for determining the primary sequence homology relationship between at least two query sequences.
  - 21. A method for determining a preferred alignment between at least one query sequence and at least one template sequence, for use in primary sequence homology modeling methods, comprising the steps of:
  - a. aligning at least two reference sequences to determine at least one
    BRIDGE/BULGE gap;
    - b. calculating a sequence alignment similarity matrix with matrix elements  $s_{ij}$  for each said query sequence and each said template sequence;

- c. calculating a sequence alignment sum matrix with matrix elements  $S_{ij}$  from the dynamic evolution of each said sequence alignment similarity matrix, according to the equation:
- $S_{ij} = s_{ij} + Max\{S_{i+1,j+1}, S_{i+1,j+2 \text{ to } jmax} GAP, S_{i+2 \text{ to } imax, j+2} GAP, S_{m.n} BRIDGE/BULGE\},$

wherein GAP represents the gap penalty for an alignment gap between said query sequence and each said template sequence, BRIDGE/BULGE represents the penalty for a known bridge or bulge that begins at the m,n matrix element of said sum matrix and ends at the i,j matrix element of said sum matrix and Max $\{S_{i+1,j+1}, S_{i+1,j+2 \text{ to jmax}} - GAP, S_{i+2 \text{ to imax}, j+2} - GAP, S_{m,n} - BRIDGE/BULGE\}$  refers to the maximum value of the four terms contained within the brackets; and

- d. determining a preferred alignment between each said query sequence and each said template sequence from the dynamic evolution of said sum matrix; wherein said preferred alignment contains from approximately 10% to approximately 20% homologous residues.
  - 22. The method of claim 21 wherein said preferred alignment is the optimal alignment.
- 20 23. The method of claim 21 wherein said primary sequence homology modeling method is a method for determining the three dimensional structure of said query sequence.

- 24. The method of claim 22 wherein said primary sequence homology modeling method is a method for determining the three dimensional structure of said query sequence.
- 25. The method of claim 21 wherein said primary sequence homology modeling
  method is a method for determining the primary sequence homology relationship
  between at least two query sequences.
  - 26. The method of claim 22 wherein said primary sequence homology modeling method is a method for determining the primary sequence homology relationship between at least two query sequences.
- 10 27 A method for determining the three dimensional structure of a query sequence based upon primary sequence homology modeling with at least one template sequence, wherein, said alignment between said query sequence and said template sequence is determined by the methods of claim 2, claim 6, claim 8, claim 12, claim 20 and claim 24.
- 28. A method for determining the primary sequence homology relationship between at least two query sequences based upon primary sequence homology modeling with at least one template sequence, wherein, said alignment between said query sequence and said template sequence is determined by the methods of claim 2, claim 6, claim 8, claim 14, claim 22, and claim 26.